PATENT COOPERATION TREATY

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INTERNATIONAL PRELIMINARY REPORT ON PATENTABILITY (Chapter II of the Patent Cooperation Treaty)

(PCT Article 36 and Rule 70)

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Applicant's or agent's file reference 913453-58PCT	FOR FURTHER ACTION	See Form PCT/IPEA/416
International application No. PCT/CA2005/000250	International filing date (day/month/year) 07 February 2005 (07-02-2005)	Priority date (day/month/year) 06 February 2004 (06-02-2004)
International Patent Classification (IPC) of IPC: C07H 21/00 (2006.01), C12Q 1	national classification and IPC /68 (2006.01), C07H 21/04 (2006.01)	
Applicant CANADIAN BLOOD SERVIC		
1. This report is the international prelimit under Article 35 and transmitted to the	ary examination report, established by this applicant according to Article 36.	International Preliminary Examining Authority
2. This REPORT consists of a total of	6 sheets, including this cover sheet.	
3. This report is also accompanied by AN	NEXES, comprising:	
	to the International Bureau) a total of	43 sheets, as follows:
[X] sheets of the des	cription, claims and/or drawings which have ntaining rectifications authorized by this Au	e been amended and are the basis of this report athority (see Rule 70.16 and Section 607 of the
[] sheets which sup goes beyond the and the Supplen	persede earlier sheets, but which this Author disclosure in the international application and application are applications.	rity considers contain an amendment that as filed, as indicated in item 4 of Box No. 1
1	Bureau only) a total of (indicate type and not only), containing a sequence listing and the Supplemental Box Relating to Sequence	number of electronic carrier(s)) d/or tables related thereto, in electronic ce Listing (see Section 802 of the Administrative
4. This report contains indications relating	ng to the following items:	
[X] Box No. I Basis of the rep	ort	
[]Box No. II Priority		
2 2	ent of opinion with regard to novelty, inven	itive step and industrial applicability
[]Box No. IV Lack of unity of		elty, inventive step or industrial applicability;
1	planations supporting such statement	city, inventive stop of industrial applications,
Box No. VI Certain docume		
[]Box No. VII Certain defects		
	tions on the international application	
Date of submission of the demand 06 December 2005 (06-	Date of completio 9 June 2006 (09-0	on of this report 06-2006)
Name and mailing address of the IPEA/C Canadian Intellectual Property Office Place du Portage I, C114 - 1st Floor, Bo 50 Victoria Street Gatineau, Quebec K1A 0C9 Facsimile No.: 001(819)953-2476	x PCT	halie Chartrand (819) 994-2341

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Box No. I	Basis of the report		
1. With	regard to the language, this report is based on:		
[X]	the international application in the language in	which it was filed	
	a translation of the international application int		, which is the language of a
	translation furnished for the purposes of:		, willou is the language of a
[] international search (Rules 12.3(a) and 2	23.1(b))	
[] publication of the international application	on (Rule 12.4(a))	
[] international preliminary examination (R	• • •	
annex	regard to the elements of the international apple ceiving Office in response to an invitation under ed to this report): he international application as originally filed/f	er Article 14 are rejerrea to in this rej	cement sheets which have been furnished in port as "originally filed" and are not
	he description:	TOT HOUSE	
[X] pages <u>1-19, 22-25, 28, 30-36 and 38.</u>	3-49	as originally filed/furnished
	X] pages* 20, 21, 21a, 26, 27, 29, 37	received by this Authority on	December 6, 2005
r	and 50-70	_	December 0, 2005
L 5327 41] pages*	received by this Authority on	
[X] tl	ne claims:	•	
l r] pages		as originally filed/furnished
L. F] pages* 71.76		ny statement) under Article 19
L	7	received by this Authority on	<u>December 6, 2005</u>
[X] th] pages* ne drawings:	received by this Authority on	
£43, ~ [pages		
ر ت			as originally filed/furnished
ָר ר		received by this Authority on	
[X] a:		received by this Authority on	
L _	sequence listing and/or any related table(s) - se	se Supplemental Box Relating to Sequ	zence Listing.
. [] Tl	ne amendments have resulted in the cancellatio	C	
[] the description, pages	M OI.	
]] the claims, Nos.		
]] the drawings, sheets/figs		
]] the sequence listing (specify):		
[] any table(s) related to sequence listing (spe	ecifv):	
	- <u>-</u>		
[] Th sin [[[[is report has been established as if (some of) the ce they have been considered to go beyond the line description, pages the claims, Nos. I the drawings, sheets/figs I the sequence listing (specify): any table(s) related to sequence listing (specify)	e disclosure as filed, as indicated in th	and listed below had not been made, see Supplemental Box (Rule 70.2(c)).
If item 4 a	applies, some or all of those sheets may be mar	rked "superseded."	

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Box No. V Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial
applicability; citations and explanations supporting such statement

applicability, citation	o torrer order		
1. Statement			
Novelty (N)	Claims	1-6 and 8-36	YES
	Claims	<u>7</u>	NO
		•	
Inventive step (IS)	Claims	1-6 and 9-36	YES
	Claims	7 and 8	NO
Industrial applicability (IA)	Claims	<u>1-36</u>	YES
	Claims	none	NO

2. Citations and explanations (Rule 70.7)

Reference is made to the documents which were cited in the written opinion of the International Searching Authority, namely:

D1: WO 01/32702 A2 (DRK BLUTSPENDEDIENST BADEN-WUERTTEMBERG GMBH) 10 May, 2001.

D2: WO 00/20634 A1 (NOVA MOLECULAR, INC.) 13 April, 2000.

D3: WO 02/068684 A2 (PYROSEQUENCING AB) 6 September, 2002.

D4: WO 02/30950 A2 (GENAISSANCE PHARMACEUTICALS, INC.) 18 April, 2002

D5: HIRSCHHORN, J. N. et al., "SBE-TAGS: An array-based method for efficient single-nucleotide polymorphism genotyping", Proceedings of the National Academy of Sciences of USA, August 2000, Vol. 97, no. 22, pages 12164-12169.

D6: GRAF, S. et al., "Genotyping of HPA-1 (Human Platelet Antigen 1) by mini-sequencing", Blood. 16 November, 2000, Vol. 96, no. 11, Part 2, page 53b.

D7: GASSNER, C. et al., "RHD/CE typing by polymerase chain reaction using sequence-specific primers", Transfusion. October 1997, Vol. 37, pages 1020-1026.

The point of invention of this application is to provide a multiplex PCR oligonucleotide extension assay to genotype a plurality of blood group or platelet antigen SNPs simultaneously.

NOVELTY AND INVENTIVE STEP under Articles 33(2) and 33(3):

D1 discloses methods to genotype RHD alleles. These methods simultaneously analyze a plurality of polymorphisms (see page 57) which comprise a step of multiplexing PCR amplification. Also, this reference discloses PCR primers used in the methods. The teaching of this reference falls within the scope of claim 7. Therefore, this claim does not comply with Article 33(2) of the PCT. In the correspondence dated December 6, 2005, the applicant argues that the teaching of D1 is different from the present application because it is restricted to a PCR methodology and primers having specificity to a single SNP of only one blood group antigen, that being RhD. However, it appears from page 57, that they analyze more than one SNP of RhD. Claim 7 does not specify that the oligonucleotide primers and probes are used to analyze a plurality of SNPs corresponding to a plurality of blood group or platelet antigen genotypes simultaneously. Therefore, the subject matter of claim 7 is encompassed by D1.

As claim 7 has been found to lack novelty under Article 33(2) of the PCT, it also lacks an inventive step under Article 33(3) of the PCT.

D3 discloses methods of allele-specific primer extension useful for detecting mutations and genetic variations. Human genomic DNA is isolated, then, multiplex PCR is performed to amplify multiple single nucleotide polymorphisms. The SNPs analyzed were wiafl 764 (A/C) on chromosome 9q, codon 72 (C/G) on the p53 gene, nucleotide position 677 (C/T) on the MTHFR gene and nucleotide position 196 (A/G) on the GPIIIa gene.

See supplemental sheet

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Box No. VIII Certain observations on the international application

The following observations on the clarity of the claims, description, and drawings or on the question whether the claims are fully supported by the description, are made:

Claim 9 is ambiguous and does not comply with Article 6 of the PCT. The preliminary paragraph of the claim defines a method of simultaneously analyzing a plurality of blood group or platelet antigens in a sample and in step b), the multiplex PCR amplification of DNA encompasses a plurality of SNPs each corresponding to <u>a</u> blood group or platelet antigen genotype. It is not clear in step b) and according to the first paragraph, whether the SNPs correspond to a plurality of blood group or platelet antigens or to a single blood group or platelet antigen.

In claim 8, the expression "more than one of all of said probe" should be replaced by "more than one or all of the said probe".

Claim 20 is indefinite and does not comply with Article 6 of the PCT. Applicant is claiming a method without fully defining it in the claim. A method is a series of steps to be followed to achieve a desired result. All of the essential steps of the allegedly novel method must be defined.

A typographic error was found in claim 20. The word "the" is repeated in the expression "more of the the oligonucleotide".

Claim 23 is indefinite and does not comply with Article 6 of the PCT. The "sample" in step (a) has no antecedent.

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Supplemental Box relating to Sequence Listing

C	ontinuatio	on of Box No.1, item 2:
1.	With reg	ard to any nucleotide and/or amino acid sequence disclosed in the international application and necessary to the claimed a, this report was established on the basis of:
	a.	type of material
		[X] a sequence listing
		[X] table(s) related to the sequence listing
	b.	format of material
		[X] on paper
		[X] in electronic form
	c.	time of filing/furnishing
		[X] contained in the international application as filed
		[] filed together with the international application in electronic form

2. [X] In addition, in the case that more than one version or copy of a sequence listing and/or table(s) relating thereto has been filed or furnished, the required statements that the information in the subsequent or additional copies is identical to that in the application as filed or does not go beyond the application as filed, as appropriate, were furnished.

December 6, 2005

furnished subsequently to this Authority for the purposes of search and/or examination

3. Additional comments:

[X] received by this Authority as an amendment* on

^{*} If item 4 in Box No. 1 applies, the listing and/or table(s) related thereto, which form part of the basis of the report, may be marked "superseded".

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Supplemental Box

In case the space in any of the preceding boxes is not sufficient.

Continuation of:

Box V

It is obvious to a skilled person, in view of D3 and common general knowledge, to prepare other primers directed to other blood group antigens in the method to identify blood group SNPs as taught in D3. Therefore, claims 7 and 8 do not define an inventive step under Article 33(3) of the PCT. As mentioned previously, claim 7, on which claim 8 depends, does not specify that the oligonucleotide primers and probes are used to analyze a plurality of SNPs corresponding to a <u>plurality</u> of blood group or platelet antigen genotypes simultaneously. Thus, claims 7 and 8 are not inventive.

The amended claims 1 to 6 and 8 to 36 submitted on December 6, 2005 appear to be novel in view of the cited documents (D1 to D7). More specifically, the applicant has amended claim 1 to specify that the nucleic acid sequences of Table 1 are for use in a PCR primer pair for multiplex SNP analysis of a plurality of blood group or platelet antigen SNPs simultaneously. The examiner agree with the argument outlined by the applicant in the correspondence dated December 6, 2005 which states that neither D1 nor D2 disclose an oligonucleotide primer and probe set for analyzing a <u>plurality</u> of blood group or platelet antigen SNPs simultaneously. Also, some claims (9, 10, 28, 29, 32 and 40) that were rejected for lack of novelty in the Written Opinion of the International Searching Authority have been deleted with the amendments submitted on December 6, 2005, thereby obviating the objection. The new claims 9, 23 and 32 now specify that the methods encompass the simultaneous analysis of a plurality of blood group or platelet antigen specific SNPs and in addition for claims 23 and 32 include the use of a plurality of primers as defined in Table 1.

Also, claims 1 to 6 and 9 to 36 appear to be inventive. The primers of claim 1 are <u>used</u> in a multiplex PCR method <u>for</u> the analysis of a plurality of blood group or platelet antigen SNPs simultaneously. Also, the method claimed in claims 9, 20, 23, 32 and the use claim 26 involve the analysis of more than one blood group or platelet antigen SNPs simultaneously. The teachings of the cited references do not describe nor suggest a methodology or primers <u>for use</u> therein for the simultaneous detection of unrelated blood group and platelet genotypes simultaneously. Therefore, the subject matter defined in those claims is novel and inventive.

INDUSTRIAL APPLICABILITY:

Claims 1 to 36 appear to have industrial applicability under Article 33(4) of the PCT, based on the use of the primers and probes of Tables 1 and 2 in a method of simultaneously analyzing a plurality of blood group or HPA antigens in a sample.